Ru-induced loss of long-range magnetic order in $a$-Fe$_{90-x}$Ru$_x$Zr$_{10}$

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Mössbauer spectroscopy, magnetization, susceptibility, and neutron depolarization have been used to investigate the loss of long-range ferromagnetic order in $a$-Fe$_{90-x}$Ru$_x$Zr$_{10}$. Our depolarization results show a rapid reduction in domain size from 1.6 to 0.4 $\mu$m as $x$ increases from 0 to 2, with no evidence for long-range order in either depolarization or magnetization data for $x \geq 3$. Furthermore, for $x=1$ and 2, we observe a clear break in the temperature dependence of $\langle B_0 \rangle$ which may be associated with ordering of transverse spin components in these partially frustrated alloys. © 1997 *American Institute of Physics*. [S0021-8979(97)17808-2]

I. INTRODUCTION

Iron-rich amorphous Fe–Zr alloys exhibit a rich variety of magnetic behavior as a result of partially frustrated exchange interaction. This frustration arises though the distance dependence of the direct exchange interaction $J(r)$ coupled with short Fe–Fe contacts in the glass which lead to significant numbers of antiferromagnetic (AF) bonds that compete with the dominant ferromagnetic (FM) interactions. Essentially all additions (e.g., hydrogen) or substitutions for Fe by other 3d transition elements or metalloids lead to an increase in ordering temperature, even when accompanied by a reduction in magnetization. The substituents act by shifting the balance between FM and AF interactions either by contributing strong FM couplings, as with Co, or by expanding the lattice, as with H, and reducing short Fe–Fe contacts. The one clear exception to this rule is Ruthenium, which leads to an immediate and rapid reduction in both $T_c$ and the magnetization when added to $a$-Fe$_{90}$Zr$_{10}$. Indeed, the only other element that comes close to the 30 K/at % effect on $T_c$ is iron itself. The reduction in $T_c$ appears to be accompanied by a loss of long-range magnetic order, and susceptibility measurements suggest that $a$-Fe$_{90-x}$Ru$_x$Zr$_{10}$ may be a spin glass (SG), with a $T_g$ of 84 K, down from a $T_c$ of 230 K in the Ru-free alloy.

We report here magnetization, Mössbauer and neutron depolarization measurements aimed at determining the nature of the Ru-induced changes in the magnetic order of $a$-Fe$_{90-x}$Ru$_x$Zr$_{10}$. Our results confirm the destruction of magnetic order and show that by $x=3$, no long-range ferromagnetic order is present above 2 K.

II. EXPERIMENTAL METHODS

The alloys were prepared by arc melting the appropriate ratio of pure elements (Fe: 99.95%, Zr: 99.5%, and Ru: 99.9% pure) under Ti-gettered argon to yield ~2 g ingots. Melt spinning was carried out under a partial pressure of Helium onto a copper wheel at 50 m/s. Absence of crystallinity was confirmed using Cu $K_\alpha$ powder x-ray diffraction and room temperature Mössbauer spectroscopy.

A LakeShore susceptibility system with a closed-cycle fridge was used to record $\chi_m$ vs $T$ down to 12 K. Magnetization data up to 7 Tesla were obtained using both a LakeShore MPMS and a Quantum design PPMS. Mössbauer measurements were made on a constant acceleration spectrometer with a 1 GBq $^{57}$CoRh source calibrated using an $\alpha$-Fe foil. Samples were mounted in a vibration-isolated closed-cycle fridge for spectra at temperatures down to 12 K.

Neutron depolarization data were obtained using the DUALSPEC triple-axis spectrometer at AECL, Chalk River. Initial polarizations of ~95% at $\lambda=0.237$ nm were achieved with $\text{Cu}_2\text{MnAl}$ single crystals as polarizer and analyzer. Measurements were made between 2 and 300 K in a 1 mT guide field, on single 5 mm lengths of 20-$\mu$m-thick ribbons.

III. RESULTS AND DISCUSSION

$\chi_m$ measurements show a steady reduction in $T_c$ and evolution from ferromagnetic behavior to a spin glass like cusp with increasing Ru content (Fig. 1). While there is an abrupt change between $x=2$ and $x=3$, the cusp for the $x=3$ sample is far more rounded than that of the $x=4$ and it is possible that some FM order is present. Magnetization at 2 K shows a decrease with increasing $x$ and an increase in high-
field slope, consistent with a loss of collinear order. These results are in complete accord with earlier work.\textsuperscript{3,4}

The temperature dependence of the average hyperfine field (derived from Window fits)\textsuperscript{5} in Fig. 2 shows an abrupt change between $x=2$ and $x=3$. For $x\leq2$, the onset of magnetic splitting exhibits conventional FM behavior and corresponds to the point where $\chi_{ac}$ rises from zero (Fig. 1), however, for $x=3$, the onset occurs at the peak in $\chi_{ac}$, as it does in spin glasses. This suggests that the SG-FM boundary may lie between $x=2$ and $x=3$. Further indications of a significant change in magnetic order at this Ru content comes from modified Arrott plots, which yield ordering temperatures consistent with our other measurements for $x\leq2$, but show no sign of order or spontaneous moment for $x>3$, strongly suggesting that this system is a spin glass for $x>2$. The ordering temperatures determined in this work are summarized in Fig. 3.

The clearest evidence for the disappearance of long-range order comes from the neutron depolarization data shown in Fig. 4. For $x=0$, we observed conventional ferromagnetic behavior: a rapid depolarization of the neutron beam on cooling through $T_c$. The curve is well fitted assuming a constant domain size, with a domain magnetization that follows a modified Brillouin function.\textsuperscript{7} Ordering temperatures for $x=0$, 1, and 2 agree with values determined by other methods. The depolarization caused at $T=0$ decreases rapidly with increasing Ru, far more so than either the magnetization or $\langle B_{hf}\rangle$ (a measure of the total iron moment), we attribute this to a rapid reduction in domain size from 1.64±0.01 $\mu$m at $x=0$ to 0.4±0.1 $\mu$m by $x=2$. For $x=3$, no depolarization is observed at any temperature, ($\Delta P<5\times10^{-4}$) and we can rule out the presence of ferromagnetic correlations on a length scale greater than 10 nm. We conclude that for $x>3$ the system is a spin glass.

For $x=1$ and 2, there is also a striking recovery in the depolarization below about 80 K. The simplest explanation for this would be a reduction in one or both of the domain size or the domain magnetization. Magnetization curves exhibit only a monotonic increase on cooling, allowing us to rule out contributions from this source, however, we have no independent measure of the domain sizes and so are unable to determine whether or not domain size changes occur. Comparison with the Mössbauer data for the $x=1$ sample raises another possibility. In the limit of small domains ($<10$ $\mu$m for these materials), the polarization goes as $P = \exp(-\alpha\lambda)^6$ where the factor $\alpha$ depends on the domain size, sample thickness, and the mean square domain magnetization perpendicular to the neutron polarization, $\langle B_{hf}^2\rangle$. For a random assembly of domains, this last term is just $2B^2/3$, while for thin sheet with its plane parallel to the neutron polarization direction and domains oriented in the plane by demagnetization fields, we get $B^2/2$. The Mössbauer spectra also contain information on spin directions through the intensity of lines 2 and 5, and for the $x=1$ sample, we observed a change in spin directions from largely perpendicular to the ribbon plane above 80 K, to more random below. If we incorporate this behavior into the calculation of $\langle B_{hf}^2\rangle$, then we are able to fit the data for the $x=1$ sample with a consistent parameter set for both the Mössbauer and depolarization data. While this consistency is suggestive, the actual domain distribution can be highly strain dependent, and differential thermal contraction can change the strain pattern on cooling. In order to unambiguously separate domain size changes from reorientations, we need to measure the depolarization.
signal for neutrons polarized along three orthogonal directions. Development of these modifications for the DUALSPEC spectrometer are currently underway.

Closer examination of $\langle B_{hy} \rangle (T)$ in Fig. 2 for $x=1, 2$ reveals a clear break in slope near 80 K. We emphasize that this break in slope is not caused by the change in the intensity of lines 2 and 5 discussed above. First, the change in intensity starts about 15 K above the break in $\langle B_{hy} \rangle (T)$. Second, forcing a constant intensity in line 2 and 5 degrades the fit quality, but does not eliminate the break in slope. The observed temperature dependence is well fitted by a combination of a modified Brillouin function and a linear term that appears only below the break temperature $T_{br}$. While this linear behavior cannot continue to $T=0$, it does give an excellent description of the data (within 0.5%). The form of the additional term was suggested by examination of the difference between the data and a Brillouin function fitted only to the high temperature ($T \gtrsim 90$ K) region. While no break is apparent in the $x=0$ data, the same Brillouin function plus linear term also fits well but yields a $T_{br}$ that is not significantly different from zero. The break points, shown in Fig. 3, lie below, but close to the ordering temperatures of the $x \geq 3$ samples. This might suggest that we are seeing two-phase behavior, with a FM component that disappears with increasing $x$ and is gradually replaced by a 80 K SG phase that dominates for $x \geq 3$. However $\langle B_{hy} \rangle (T)$ is not linear in $T$ for $x \geq 3$, rather it follows the same modified Brillouin function found for $x<3$. Furthermore, we detect no indication of two-phase behavior in other measurements: (i) Magnetization curves show no change in form on heating through the break temperature. (ii) The width of the hyperfine field distribution increases gradually on cooling and does not exhibit the sharpening expected when an additional component orders. (iii) $\langle B_{hy} \rangle (0 \text{ K})$ does not change dramatically between $x=2$ and 3, the difference is only 1 T, even though the linear term, which we might associate with the SG phase, never contributes more than 5 T and should be the only component present at $x=3$.

An alternative, uniform, description of the system is provided by bond frustration. Increasing numbers of frustrated bonds lead to reduced $T_c$ and magnetization, and to the appearance of a second transition ($T_{xy}$) below which noncollinear order develops. $T_c$ and $T_{xy}$ merge at high frustration levels and the system becomes a spin glass. Such a picture has been used to account for the magnetic properties of $\alpha$-Fe$_{86}$Zr$_{14}$ and is fully consistent with the phase diagram in Fig. 3 if $T_{br}$ is identified with $T_{xy}$. The only remaining question is why Ru additions lead to an increase in exchange frustration. A simple appeal to atomic packing fails because Ru is larger than Fe. The increase in the numbers of short Fe–Fe distances that follow from replacing a large Zr atom by a smaller Fe has been used to explain increasing frustration in a $\alpha$-Fe$_{86}$Zr$_{14}$, however, for Ru such an argument would predict a reduction in frustration. Similarly, we can neither appeal to antiferromagnetically couple Ru moments as isolated Ru atoms do not carry a moment in Fe, nor would such isolated AF moments cause the frustration needed to drive the system towards spin glass ordering. One possible mechanism might be that the changes in the density of states associated with Ru additions act so as to change the crossover point in $\mathcal{T}(r)$ so that the distance at which the Fe–Fe exchange interaction switches from Fm to AF is increased and more bonds become frustrated.

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